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(\$4) Title: METHODS OF OPTIMIZING ANTIBODY VARIABLE REGION BINDING AFFINITY

(57) Abstract: The invention provides a method of conferring donor CDR binding affinity onto an antibody acceptor variable region framework. The invention also provides a method of simultaneously grafting and optimizing the binding affinity of a variable region binding fragment. A method of optimizing the binding affinity of an antibody variable region is also provided. The variable region populations can be heavy or light chains and can be expressed as individual populations or they can be coexpressed to produce heteromeric variable region binding fragments.

1

# Methods of Optimizing Antibody Variable Region Binding Affinity

# BACKGROUND OF THE INVENTION

This invention relates generally to a method of monoclonal antibody production and specifically to the simultaneous *in vitro* affinity optimization of multiple distinct domains of a variable region of a monoclonal antibody.

The War on Cancer is entering its third decade

10 and recent years have shown tremendous progress in the

understanding of cancer development and progression yet

there has been only marginal decreases in death rates

from most types of cancer. Standard chemotherapy and

radiation therapy generally involve treatment with

15 therapeutic agents that impact not only cancer cells but

other highly proliferative cells of the body, often

leading to debilitating side effects. Thus, it is

desirable to identify therapeutic agents with a higher

degree of specificity for the carcinogenic lesion.

The discovery of monoclonal antibodies (mAbs) in the 1970's provided great hope for the reality of creating therapeutic molecules with high specificity. Antibodies that bind to tumor antigens would provide specific targeting agents for cancer therapy. However, while the development of monoclonal antibodies has provided a valuable diagnostic reagent, certain limitations restrict their use as therapeutic entities.

A limitation encountered when attempts are made to use mAbs as therapeutic agents is that since mAbs are

2

developed in non-human species, usually mouse, they elicit an immune response in human patients. Chimeric antibodies join the variable region of the non-human species, which confers binding activity, to a human constant region. However, the chimeric antibody is often still immunogenic and it is therefore necessary to further modify the variable region.

One modification is the grafting of complementarity-determining regions, (CDRs) which are in 10 part antigen binding onto a human antibody variable framework. However, this approach is imperfect because CDR grafting often diminishes the binding activity of the resulting humanized mAb. Attempts to regain binding activity require laborious, step-wise procedures which 15 have been pursued essentially by a trial and error type of approach. For example, one difficulty in regaining binding affinity is because it is difficult to predict which framework residues serve a critical role in maintaining antigen binding affinity and specificity. 20 Consequently, while antibody humanization methods that rely on structural and homology data are used, the complexity that arises from the large number of framework residues potentially involved in binding activity has prevented success.

Combinatorial methods have been applied to restore binding affinity, however, these methods require sequential rounds of mutagenesis and affinity selection that can both be laborious and unpredictable.

Thus, there exists a need for efficient and reliable methods for producing human monoclonal antibodies which exhibit comparable or enhanced binding affinities to their non-human counterparts. The present

3

invention satisfies this need and provides related advantages as well.

#### SUMMARY OF THE INVENTION

The invention provides a method of conferring 5 donor CDR binding affinity onto an antibody acceptor variable region framework. The method consists of: (a) constructing a population of altered antibody variable region encoding nucleic acids, said population comprising encoding nucleic acids for an acceptor 10 variable region framework containing a plurality of different amino acids at one or more acceptor framework region amino acid positions and donor CDRs containing a plurality of different amino acids at one or more donor CDR amino acid positions; (b) expressing said population 15 of altered variable region encoding nucleic acids, and (c) identifying one or more altered variable regions having binding affinity substantially the same or greater than the donor CDR variable region. The acceptor variable region framework can be a heavy or light chain 20 variable region framework and the populations of heavy and light chain altered variable regions can be expressed alone to identify heavy or light chains having binding affinity substantially the same or greater than the donor CDR variable region. The populations of heavy and light 25 chains additionally can be coexpressed to identify heteromeric altered variable region binding fragments. The invention also provides a method of simultaneously grafting and optimizing the binding affinity of a variable region binding fragment. The method consists 30 of: (a) constructing a population of altered heavy chain variable region encoding nucleic acids comprising an acceptor variable region framework containing donor CDRs and a plurality of different amino acids at one or more

4

PCT/US00/28435

framework region and CDR amino acid positions; (b) constructing a population of altered light chain variable region encoding nucleic acids comprising an acceptor variable region framework containing donor CDRs 5 and a plurality of different amino acids at one or more framework regions and CDR amino acid positions; (c) coexpressing said populations of heavy and light chain variable region encoding nucleic acids to produce diverse combinations of heteromeric variable region 10 binding fragments, and (d) identifying one or more heteromeric variable region binding fragments having affinity substantially the same or greater than the donor CDR heteromeric variable region binding fragment. A method of optimizing the binding affinity of an antibody 15 variable region is also provided. The method consists of: (a) constructing a population of antibody variable region encoding nucleic acids, said population comprising two or more CDRs containing a plurality of different amino acids at one or more CDR amino acid positions; 20 (b) expressing said population of variable region encoding nucleic acids, and (c) identifying one or more variable regions having binding affinity substantially the same or greater than the donor CDR variable region. The variable region populations can be heavy or light 25 chains and can be expressed as individual populations or they can be coexpressed to produce heteromeric variable region binding fragments.

### BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows the alignment of anti-CD40 30 variable region and human template amino acid sequences.

Figure 2 shows binding reactivity of humanized anti-CD40 variants.

5

Figure 3 shows molecular modeling of anti-CD40 variant CW43.

Figure 4 shows a comparison of the quantitation of murine framework residues in active variants from two 5 libraries.

#### DETAILED DESCRIPTION OF THE INVENTION

The invention is directed to a method of conferring donor CDR binding affinity onto an antibody 10 acceptor variable region framework. The method effectively combines CDR grafting procedures and affinity reacquisition of the grafted variable region into a single step. The methods of the invention also are applicable for affinity maturation of an antibody 15 variable region. The affinity maturation process can be substituted for, or combined with the affinity reacquisition function when being performed during a CDR grafting procedure. Alternatively, the affinity maturation procedure can be performed independently from 20 CDR grafting procedures to optimize the binding affinity of variable region, or an antibody. An advantage of combining grafting and affinity reacquisition procedures, or affinity maturation, is the avoidance of time consuming, step-wise procedures to generate a grafted 25 variable region, or antibody, which retains sufficient binding affinity for therapeutic utility. Therefore, therapeutic antibodies can be generated rapidly and efficiently using the methods of the invention. Such advantages beneficially increase the availability and 30 choice of useful therapeutics for human diseases as well as decrease the cost to the developer and ultimately to the consumer.

PCT/US00/28435

In one embodiment, the invention is directed to methods of producing grafted heavy and light chain. variable regions having similar or better binding affinity as the CDR donor variable region. When 5 coexpressed, the grafted heavy and light chain variable regions assemble into variable region binding fragments having similar or better binding affinity as the donor antibody or variable region binding fragments thereof. The grafting is accomplished by generating a diverse 10 library of CDR grafted variable region fragments and then screening the library for binding activity similar or better than the binding activity of the donor. A diverse library is generated by selecting acceptor framework positions that differ at the corresponding position 15 compared to the donor framework and making a library population containing of all possible amino acid residue changes at each of those positions together with all possible amino acid residue changes at each position within the CDRs of the variable region. The grafting is 20 accomplished by splicing a population of encoding nucleic acids for the donor CDR containing species representing all possible amino acid residues at each CDR position into a population of encoding nucleic acids for an antibody acceptor variable region framework which 25 contains species representing all possible amino acid residue changes at the selected framework positions. The resultant population encodes the authentic donor and acceptor framework amino acid sequences as well as all possible combinations and permutations of these sequences 30 with each of the 20 naturally occurring amino acids at the changed positions.

In another embodiment, the invention is directed to methods of producing grafted heavy and light chain variable regions, and heteromeric binding fragments

7

thereof, having similar or better binding affinity as the CDR donor variable region. As described above, the grafting is accomplished by generating a diverse library of CDR grafted variable region fragments and then 5 screening the library for binding activity similar or better than the binding activity of the donor. However, the diverse library is generated by selecting acceptor framework positions that are predicted to affect CDR binding affinity and making a library population 10 containing of all possible amino acid residue changes at each of those positions or subsets of the selected amino acid positions together with all possible amino acid residue changes at each position within the CDRs of the variable region, or subsets of CDR positions. The 15 grafting is accomplished by splicing a population of encoding nucleic acids for the donor CDR containing the selected position changes into a population of encoding nucleic acids for an antibody acceptor variable region framework which contains the selected position changes.

In yet another embodiment, the invention is directed to the optimization of binding affinity of an antibody variable region. The optimization is accomplished by generating a library of variable regions which contain all possible amino acid residue changes at each amino acid position within two or more CDRs. When expressed and screened for binding activity, the variable region, or heavy and light chain heteromeric binding fragments, those species within the population are selected that contain increased or decreased binding activity compared to the parent molecule as optimal binders. Libraries containing subsets, representing less than all amino acid positions within the CDRs, can similarly be generated and screened for selecting optimal

binding variable regions and heteromeric binding fragments thereof.

As used herein, the term "CDR" or "complementarity determining region" is intended to mean 5 the non-contiguous antigen combining sites found within the variable region of both heavy and light chain polypeptides. These particular regions have been described by Kabat et al., J. Biol. Chem. 252, 6609-6616 (1977) and Kabat et al., Sequences of protein of 10 immunological interest. (1991), and by Chothia et al., J. Mol. Biol. 196:901-917 (1987) and by MacCallum et al., <u>J. Mol. Biol.</u> 262:732-745 (1996) where the definitions include overlapping or subsets of amino acid residues when compared against each other. Nevertheless, 15 application of either definition to refer to a CDR of an antibody or grafted antibodies or variants thereof is intended to be within the scope of the term as defined and used herein. The amino acid residues which encompass the CDRs as defined by each of the above cited references 20 are set forth below in Table 1 as a comparison.

Table 1: CDR Definitions

		<u>Kabat<sup>1</sup></u>	<u>Chothia<sup>2</sup></u>	MacCallum <sup>3</sup>
	V <sub>H</sub> CDR1	31-35	26-32	30-35
	VH CDR2	50-65	53-55	47-58
25	V <sub>H</sub> CDR3	95-102	96-101	93-101
	V <sub>L</sub> CDR1	24-34	26-32	30-36
	V <sub>L</sub> CDR2	50-56	50-52	46-55
	V <sub>L</sub> CDR3	89-97	91-96	89-96

Residue numbering follows the nomenclature of Kabat et 30 al., supra

<sup>&</sup>lt;sup>2</sup> Residue numbering follows the nomenclature of Chothia et al., supra

9

<sup>3</sup> Residue numbering follows the nomenclature of MacCallum et al., supra

As used herein, the term "framework" when used in reference to an antibody variable region is entered to 5 mean all amino acid residues outside the CDR regions within the variable region of an antibody. Therefore, a variable region framework is between about 100-120 amino acids in length but is intended to reference only those amino acids outside of the CDRs.

10 As used herein, the term "framework region" is intended to mean each domain of the framework that is separated by the CDRs. Therefore, for the specific example of a heavy chain variable region and for the CDRs as defined by Kabat et al., framework region 1 15 corresponds to the domain of the variable region encompassing amino acids 1-30; region 2 corresponds to the domain of the variable region encompassing amino acids 36-49; region 3 corresponds to the domain of the variable region encompassing amino acids 66-94, and 20 region 4 corresponds to the domain of the variable region from amino acids 103 to the end of the variable region. The framework regions for the light chain are similarly separated by each of the light claim variable region CDRs. Similarly, using the definition of CDRs by Chothia 25 et al. or McCallum et al. the framework region boundaries are separated by the respective CDR termini as described above.

As used herein, the term "donor" is intended to
30 mean a parent antibody molecule or fragment thereof from
which a portion is derived from, given or contributes to
another antibody molecule or fragment thereof so as to
confer either a structural or functional characteristic

10

of the parent molecule onto the receiving molecule. For the specific example of CDR grafting, the parent molecule from which the grafted CDRs are derived is a donor molecule. The donor CDRs confer binding affinity of the parent molecule onto the receiving molecule. It should be understood that a donor molecule does not have to be from a different species as the receiving molecule of fragment thereof. Instead, it is sufficient that the donor is a separate and distinct molecule.

As used herein, the term "acceptor" is intended to mean an antibody molecule or fragment thereof which is to receive the donated portion from the parent or donor antibody molecule or fragment thereof. An acceptor antibody molecule or fragment thereof is therefore

15 imparted with the structural or functional characteristic of the donated portion of the parent molecule. For the specific example of CDR grafting, the receiving molecule for which the CDRs are grafted is an acceptor molecule. The acceptor antibody molecule or fragment is imparted

20 with the binding affinity of the donor CDRs or parent molecule. As with a donor molecule, it is understood that an acceptor molecule does not have to be from a different species as the donor.

A "variable region" when used in reference to
25 an antibody or a heavy or light chain thereof is intended
to mean the amino terminal portion of an antibody which
confers antigen binding onto the molecule and which is
not the constant region. The term is intended to include
functional fragments thereof which maintain some of all
30 of the binding function of the whole variable region.
Therefore, the term "heteromeric variable region binding
fragments" is intended to mean at least one heavy chain
variable region and at least one light chain variable

11

WO 01/27160 PCT/US00/28435

regions or functional fragments thereof assembled into a heteromeric complex. Heteromeric variable region binding fragments include, for example, functional fragments such as Fab, F(ab)2, Fv, single chain Fv (scFv) and the like. 5 Such functional fragments are well known to those skilled in the art. Accordingly, the use of these terms in describing functional fragments of a heteromeric variable region is intended to correspond to the definitions well known to those skilled in the art. Such terms are 10 described in, for example, Harlow and Lane, Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory, New York (1989); Molec. Biology and Biotechnology: A Comprehensive Desk Reference (Myers, R.A. (ed.), New York: VCH Publisher, Inc.); Huston et al., Cell 15 Biophysics, 22:189-224 (1993); Plückthun and Skerra, Meth. Enzymol., 178:497-515 (1989) and in Day, E.D., Advanced Immunochemistry, Second Ed., Wiley-Liss, Inc., New York, NY (1990).

20 As used herein, the term "population" is intended to refer to a group of two or more different molecules. A population can be as large as the number of individual molecules currently available to the user or able to be made by one skilled in the art. Populations 25 can be as small as 2-4 molecules or as large as  $10^{13}$ molecules. An example where a small population can be useful is where one wishes to optimize binding affinity of a variable region or of heteromeric binding fragments by compiling beneficial differences from a small number 30 of parent molecules having similar binding affinity into a single variable binding fragment species. An example of where large populations, including as large as 108 or greater different molecules, can be desired is where all possible combinations of amino acids differences between 35 donor and acceptor at all positions within a variable

12

region are to be generated in order to obtain maximum diversity and increase the efficiency of compiling beneficial changes. In some embodiments, populations are between about 5 and 10 different species as well as up to hundreds or thousands of different species. The populations can be diverse or redundant depending on the intent and needs of the user. Those skilled in the art will know what size and diversity of a population is suitable for a particular application.

10 As used herein, the term "altered" when used in reference to an antibody variable region is intended to mean a heavy or light chain variable region that contains one or more amino acid changes in a framework region, a CDR or both compared to the parent amino acid sequence at 15 the changed position. Where an altered variable region is derived from or composed of different donor and acceptor regions, the changed amino acid residues within the altered species are to be compared to their respective amino acid positions within the parent donor 20 and acceptor regions. For example, a variable region containing donor CDRs grafted into an acceptor framework and containing one or more amino acid changes within the framework regions and one or more amino acid changes within the CDRs will have amino acids residues at the 25 changed framework region positions different than the residues at the comparable positions in the acceptor framework. Similarly, such an altered variable region will have amino acid residues at the changed CDR positions different than the residues at the comparable 30 positions in the donor CDRs.

As used herein, the term "nucleic acid" or "nucleic acids" is intended to mean a single- or double-stranded DNA or RNA molecule. A nucleic acid molecule of

13

the invention can be of linear, circular or branched configuration, and can represent either the sense or antisense strand, or both, of a native nucleic acid molecule. The term also is intended to include nucleic 5 acid molecules of both synthetic and natural origin. A nucleic acid molecule of natural origin can be derived from any animal, such as a human, non-human primate, mouse, rat, rabbit, bovine, porcine, ovine, canine, feline, or amphibian, or from a lower eukaryote, such as 10 Drosophila, C. elegans or yeast. A synthetic nucleic acid includes, for example, chemical and enzymatic synthesis. The term "nucleic acid" or "nucleic acids" is similarly intended to include analogues of natural nucleotides which have similar functional properties as 15 the referenced nucleic acid and which can be utilized in a manner similar to naturally occurring nucleotides and nucleosides.

As used herein, the term "coexpressing" is intended to mean the expression of two or more molecules 20 by the same cell. The coexpressed molecules can be polypeptides or encoding nucleic acids. The coexpression can be, for example, constitutive or inducible. Such nucleic acid sequences can also be expressed simultaneously or, alternatively, regulated 25 independently. Various combinations of these modes of coexpression can additionally be used depending on the number and intended use of the variable region encoding nucleic acids. The term is intended to include the coexpression of members originating from different 30 populations in the same cell. For example, populations of molecules can be coexpressed where single or multiple different species from two or more populations are expressed in the same cell. A specific example includes the coexpression of heavy and light chain variable region

14

PCT/US00/28435

populations where at least one member from each population is expressed together in the same cell to produce a library of cells coexpressing different species of heteromers variable region binding fragments.

of heteromers variable region binding fragments.

Populations which can be coexpressed can be as small as 2 different species within each population. Additionally, the number of molecules coexpressed from different populations also can be as large as 10° or greater, such as in the case where multiple amino acid position changes of multiple framework regions or CDRs in both heavy and light chain antibody variable region populations are produced and coexpressed. Numerous different sized populations of encoding nucleic acids inbetween the the above ranges and greater can also be coexpressed. Those skilled in the art know, or can determine, what modes of coexpression can be used to achieve a particular goal or satisfy a desired need.

As used herein, the term "identifying" is intended to mean detecting by a qualitative or 20 quantitative means, a variable region or altered variable of the invention by functional or biochemical properties, including, for example, binding affinity of catalytic activity.

As used herein the term "binding affinity" is intended to mean the strength of a binding interaction and therefore includes both the actual binding affinity as well as the apparent binding affinity. The actual binding affinity is a ratio of the association rate over the disassociation rate. Therefore, conferring or optimizing binding affinity includes altering either or both of these components to achieve the desired level of binding affinity. The apparent affinity can include, for example, the avidity of the interaction. For example, a

15

bivalent heteromeric variable region binding fragment can exhibit altered or optimized binding affinity due to its valency.

As used herein, the term "optimizing" when used 5 in reference to a variable region or a functional fragment thereof is intended to mean that the binding affinity of the variable region has been modified compared to the binding affinity of a parent variable region or a donor variable region. A variable region 10 exhibiting optimized activity can exhibit, for example, higher affinity or lower affinity binding, or increased or decreased association or dissociation rates compared to an unaltered variable region. A variable region exhibiting optimized activity also can exhibit increased 15 stability such as increased half-life in a particular organism. For example, an antibody activity can be optimized to increase stability by decreasing susceptibility to proteolysis. An antibody exhibiting optimized activity also can exhibit lower affinity 20 binding, including decreased association rates or increased dissociation rates, if desired. An optimized variable region exhibiting lower affinity binding is useful, for example, for penetrating a solid tumor. In contrast to a higher affinity variable region, which 25 would bind to the peripheral regions of the tumor but would be unable to penetrate to the inner regions of the tumor due to its high affinity, a lower affinity variable region would be advantageous for penetrating the inner regions of the tumor. As with optimization of binding 30 affinities above, optimization of a catalytic variable region can be, for example, increased or decreased catalytic rates, disassociation constants or association constants.

16

PCT/US00/28435

As used herein, the term "substantially the same" when used in reference to binding affinity is intended to mean similar or identical binding affinities where one molecule has a binding affinity constant that is similar to another molecule within the experimental variability of the affinity measurement. The experimental variability of the binding affinity measurement is dependent upon the specific assay used and is known to those skilled in the art.

The invention provides a method for conferring 10 donor CDR binding affinity onto an antibody acceptor variable region framework. The method consists of: (a) constructing a population of altered antibody variable region encoding nucleic acids, the population consisting 15 of encoding nucleic acids for an acceptor variable region framework containing a plurality of different amino acids at one or more acceptor framework region amino acid positions and donor CDRs containing a plurality of different amino acids at one or more donor CDR amino acid 20 positions; (b) expressing the population of altered variable region encoding nucleic acids, and (c) identifying one or more altered variable regions having binding affinity substantially the same or greater than the donor CDR variable region.

The process of producing human antibody forms from nonhuman species involves recombinantly splicing CDRs from a nonhuman donor antibody into a human acceptor framework region to confer binding activity onto the resultant grafted antibody, or variable region binding fragment thereof. The process of grafting, referred to as the procedure for splicing CDRs into a framework, while mechanically simple it almost always results in a grafted antibody that exhibits a substantial loss in

17

PCT/US00/28435

binding affinity. Although donor and acceptor variable regions are structurally similar, the process nevertheless combines CDR binding domains with a heterologous acceptor region, resulting in a 5 conformationally imperfect setting for the binding residues of the grafted antibody. Therefore, once the CDR-grafted antibody, or variable region binding fragment is made, it requires subsequent rounds of molecular engineering to reacquire binding affinity comparable to 10 the donor antibody. The present invention combines these steps such that CDR grafting and binding reacquisition occur in a single simultaneous procedure. The method is also applicable to optimizing the binding affinity of an antibody, or variable region binding fragment 15 simultaneous with CDR grafting and to optimizing an antibody or variable region binding fragment in a single procedure without including the CDR grafting process.

The methods of the invention confer or impart donor CDR binding affinity onto an antibody acceptor

variable region framework in a procedure which achieves grafting of donor CDRs and affinity reacquisition in a simultaneous process. The methods similarly can be used, either alone or in combination with CDR grafting, to modify or optimize the binding affinity of a variable region. The methods for conferring donor CDR binding affinity onto an acceptor variable region are applicable to both heavy and light chain variable regions and as such can be used to simultaneous graft and optimize the binding affinity of an antibody variable region.

30 The methods for conferring donor CDR binding affinity onto a variable region involve identifying the relevant amino acid positions in the acceptor framework that are known or predicted to influence a CDR

18

PCT/US00/28435

conformation, or that are known or predicted to influence the spacial context of amino acid side chains within the CDR that participate in binding, and then generate a population of altered variable region species that 5 incorporate a plurality of different amino acid residues at those positions. For example, the different amino acid residues at those positions can be incorporated either randomly or with a predetermined bias and can include all of the twenty naturally occurring amino acid 10 residues at each of the relevant positions. Subsets, including less than all of the naturally occurring amino acids can additionally be chosen for incorporation at the relevant framework positions. Including a plurality of different amino acid residues at each of the relevant 15 framework positions ensures that there will be at least one species within the population that will have framework changes which allows the CDRs to reacquire their donor binding affinity in the context of the acceptor framework variable region.

selected amino acid positions, the CDRs also are altered to contain a plurality of different amino acid residue changes at all or selected positions within the donor CDRs. For example, random or biased incorporation of the twenty naturally occurring amino acid residues, or preselected subsets, are also introduced into the donor CDRs to produce a diverse population of CDR species. Including a diverse population of different CDR variant species ensures that beneficial changes in the framework positions are not neutralized by a conformationally incompatible residue in a donor CDR. Inclusion of CDR variant species into the diverse population of variable regions also allows for the generation of variant species

19

that exhibit optimized binding affinity for a predetermined antigen.

The resultant population of CDR grafted variable regions described above will therefore contain,

5 at the relevant framework positions and at the selected CDR positions, a species corresponding to the authentic parent amino acid residue at each position as well as a diverse number of different species which correspond to the possible combinations and permutations of the

10 authentic parent amino acid residues together with the variant residues at each of the relevant framework and selected CDR positions. Such a diverse population of CDR grafted variable regions are screened for an altered variable region species which retains donor CDR binding

15 activity, or optimized binding activity.

One advantage of the methods of the invention is that they do not limit the choice of acceptor variable regions applicable, or expected to be successful, for receiving CDRs from the donor molecule. For example, 20 when choosing an acceptor region it can be desirable, or in some circumstances even required, to select an acceptor that is closely similar to the variable region amino acid sequence harboring the donor CDRs because the CDR conformation in the grafted variable region will 25 likely be more similar to that of the donor. However, selecting similar framework region sequences between the donor and acceptor variable regions still does not provide which residues, out of the differences, actually play a role in CDR binding affinity of the grafted 30 variable region. Selection of similar acceptor frameworks therefore only limits the number of possible residues which to investigate in order to reacquire binding affinity onto the grafted variable region. The

20

methods of the invention circumvent this problem by producing a library of all possible or relevant changes in the acceptor framework, and then screening those variable regions, or heteromeric binding fragments

5 thereof for species that maintain or exhibit increased binging affinity compared to the donor molecule.

Therefore, the applicability is not preconditioned on the availability or search for an acceptor framework variable region similar to that of the donor.

Selection of the relevant framework amino acid 10 positions to alter can depend on a variety of criteria well known to those skilled it the art. As described above, one criteria for selecting relevant framework amino acids to change can be the relative differences in 15 amino acid framework residues between the donor and acceptor molecules. Selection of relevant framework positions to alter using this approach is simple and has the advantage of avoiding any subjective bias in residue determination or any inherent bias in CDR binding 20 affinity contribution by the residue. Criteria other than relatedness of amino acid residues can be used for selecting relevant framework positions to alter. criteria can be used in combination with, or alternative to the selection of framework positions having divergent 25 amino acid residues. These additional criteria are described further and similarly are well known to those skilled in the art.

Another criteria which can be used for determining the relevant amino acid positions to change 30 can be, for example, selection of framework residues that are known to be important, or contribute to CDR conformation. For example, canonical framework residues play such a role in CDR conformation or structure. Such

21

residues can be considered to be relevant to change for a variety of reasons, including for example, their new context of being associated with heterologous CDR sequences in the grafted variable region. Targeting of a canonical framework residue as a relevant position to change can identify a more compatible amino acid residue in context with its associated donor CDR sequence. Additionally, targeting of canonical residues can allow for the identification of residues at these positions that absorb detrimental effects to CDR structure from residues located elsewhere in the framework region.

The frequency of an amino acid residue at a particular framework position is another criteria which can be used for selecting relevant framework amino acid positions to change. For example, comparison of the selected framework with other framework sequences within its subfamily can reveal residues that occur at minor frequences at a particular position or positions. Such positions harboring less abundant residues are similarly applicable for selection as a position to alter in the acceptor variable region framework.

The relevant amino acid positions to change also can be selected, for example, based on proximity to a CDR. In certain contexts, such residues can participate in CDR conformation or antigen binding. Moreover, this criteria can similarly be used to prioritize relevant positions selected by other criteria described herein. Therefore, differentiating between residues proximal and distal to one or more CDRs is an efficient way to reduce the number of relevant positions to change using the methods of the invention.

22

Other criteria for selecting relevant amino acid framework positions to alter include, for example, residues that are known or predicted to reside in threedimensional space near the antigen-CDR interface or 5 predicted to modulate CDR activity. Similarly, framework residues that are known or predicted to contact opposite domain of the heavy  $(V_u)$  and light  $(V_t)$  chain variable region interface. Such framework positions can effect the conformation or affinity of a CDR by modulating the 10 CDR binding pocket, antigen interaction or the  $V_{\rm H}$  and  $V_{\rm L}$ interaction. Therefore, selection of these amino acid positions as relevant for construction of the diverse population to screen can beneficially identify framework changes which replace residues having detrimental effects 15 on CDR conformation or absorb detrimental effects of residues occurring elsewhere in the framework.

Finally, other framework residues that can be selected for alteration include amino acid positions that are inaccessible to solvent. Such residues are generally buried in the variable region and therefore capable of influencing the conformation of the CDR or V<sub>H</sub> and V<sub>L</sub> interactions. Solvent accessibility can be predicted, for example, from the relative hydrophobicity of the environment created by the amino acid side chains of the polypeptide or by known three-dimensional structural data.

In addition to selecting the relevant framework positions, the method of conferring donor CDR binding affinity onto an antibody acceptor variable region

framework also incorporates changes in the donor CDR amino acid positions. As with selecting the relevant framework positions to change, there is similarly a range of possible changes that can be made in the donor CDR

23

positions. Some or all of the possible changes that can be selected for change can be introduced into the population of grafted donor CDRs to practice the methods of the invention.

5 One approach is to change all amino acid positions along a CDR by replacement at each position with, for example, all twenty naturally occurring amino acids. The replacement of each position can occur in the context of other donor CDR amino acid positions so that a 10 significant portion of the CDR maintains the authentic donor CDR sequence, and therefore, the binding affinity of the donor CDR. For example, an acceptor variable region framework targeted for relevant amino acid positions changes as described previously, can be 15 targeted for grafting with a population of CDRs containing single position replacements at each position within the CDRs. Similarly, an acceptor variable region framework can be targeted for grafting with a population of CDRs containing more than one position changed to 20 incorporate all twenty amino acid residues, or a fractional subset, at each set of positions within the CDRs. For example, all possible sets of changes corresponding to two, three or four or more amino acid positions within a CDR can be targeted for introduction 25 into a population of CDRs for grafting into an acceptor variable region framework.

Single amino acid position changes are generated at each position without altering the remain amino acid positions within the CDR. A population of single position changes will contain at each position the varied amino acid residues, incorporated either randomly or with a biased frequency, while leaving the remaining positions as donor CDR residues. For the specific

example of a ten residue CDR, the population will contain species having the first, second and third, continued through the tenth CDR residue, individually randomized or represented by a biased frequency of incorporated amino acid residues while the remaining non-varied positions represent the donor CDR amino acid residues. For the specific example described above, these non-varied positions would correspond to positions 2-10; 1,3-10; 1,2,4-10, continued through positions 1-9, respectively.

Therefore, the resultant population will contain species that represent all single position changes.

24

Similarly, double, triple and quadruple amino acid position changes can be generated for each set of positions without altering the remain amino acid 15 positions within the CDR. For example, a population of double position changes will contain at each set of two positions the varied amino acid residues while leaving the remaining positions as donor CDR residues. The sets will correspond to, for example, positions 1 and 2, 1 and 20 3, 1 and 4, through the set corresponding to the first and last position of the CDR. The population will also contain sets corresponding to positions 2 and 3, 2 and 4, 2 and 5, through the set corresponding to the second an last position of the CDR. Likewise, the population will 25 contain sets of double position changes corresponding to all pairs of position changes beginning with position three of the CDR. Similar pairs of position changes are made with the remaining sets CDR amino acid positions. Therefore, the population will contain species that 30 represent all pairwise combinations of amino acid position changes. In a simialar fashion, populations corresponding to sets of changes representing all triple and quadruplet changes along a CDR can similarly be

25

targeted for grafting into the variable region frameworks using the methods of the invention.

The above populations of CDR variant species can be targeted for any or all of the CDRs which 5 constitute the binding pocket of a variable region. Therefore, an acceptor variable region framework targeted for relevant amino acid positions changes as described previously, can be targeted for the simultaneous incorporation of donor CDR variant populations at one, 10 two or all three recipient CDR locations. The choice of which CDR or the number of CDRs to target with amino acid position changes will depend on, for example, if a full CDR grafting into an acceptor is desired or whether the method is being performed for optimization of binding 15 affinity. Many grafting procedures will generally employ the grafting of all three CDRs, where at least one of the CDRs will contain amino acid positions changes. Generally however, all of the donor CDRs will be populations containing amino acid position changes. 20 Converesly, and as described further below, optimization procedures can employ CDR variant populations corresponding to any or all of the CDRs within a variable region.

Another approach for selecting donor CDR amino
25 acids to change for conferring donor CDR binding affinity
onto an antibody acceptor variable region framework is to
select known or readily identifiable CDR positions that
are highly variable. For example, the variable region
CDR 3 is generally highly variable due to genetic
30 recombination. This region therefore can be selectively
targeted for amino acid position changes during grafting
procedures to ensure binding affinity reacquisition or

26

augmentation when made together with relevant acceptorvariable framework changes as described previously.

In contrast, CDR residues that appear conserved or have been empirically determined to be non-mutable by 5 functional criteria will generally be avoided when selecting residues in the CDR to target for change. It should be noted however, that apparent non-mutable residues can nevertheless be successfully changed using the methods of the invention because the populations of 10 altered variable regions contain from a few to many amino acid position changes in both the framework regions and in the CDR regions. As such, the CDR grafted variable regions identified by binding affinity are a result of the all the changes and therefore, all the interactions 15 of residues introduced into a particular species. Therefore, suboptimal residues incorporated at, for example, an apparent non-mutable position can be counteracted and even augmented by amino acid substitutions elsewhere in the framework regions or in 20 other CDRs.

Similarly, because the methods of the invention for CDR grafting, affinity reacquisition and affinity optimization employ the production and screening of diverse populations of variable region species generated from an acceptor framework and donor CDR variants, there are numerous effects on binding affinity that will occur due to the combined interactions of two or more amino acid changes within a single variable region species. For example, the affect of amino acid changes in either a framework region or CDR that are inherently beneficial can be masked or neutralized due to surrounding authentic parent residues or due to their context in a heterologous region of a grafted antibody. However, second site

changes in the surrounding residues or the heterologous regions can unveil the beneficial characteristics of the latent residue or residues. Such second site changes can occur, for example, in both proximal and distal heterologous or homologous region sequences.

27

For example, if the beneficial residue is in a grafted CDR region, the proximal heterologous sequences would be the adjacent framework regions whereas distal heterologous regions would be framework regions separated 10 by an adjacent CDR. In this specific example, a proximal homologous region would be the surrounding residues within the grafted CDR harboring the beneficial change whereas the remaining CDRs are examples of distal homologous regions. By analogy, the opposite would be 15 true for a inherently beneficial residue in a framework region. Specifically, proximal homologous region sequences would be located in the same framework region and distal homologous sequences would be in any of the other framework regions. Proximal heterologous region 20 sequences would be in the adjacent CDR or CDRs whereas nonadjacent CDRs constitute distal heterologous region sequences. Such second site effects can occur, for example, through the translation of conformational changes to the CDR binding pocket or to the framework 25 regions.

Other effects on binding affinity that will occur due to the combined interactions of two or more amino acid changes within a single variable region species include, for example, the neutralization or augmentation of inherently detrimental changes and the augmentation of beneficial amino acid changes or the augmentation of parent residues. As with the unveiling of beneficial changes and the ability to counteract

28

changes in apparently non-mutable residues, the neutralization and augmentation of amino acid changes within the grafted CDRs or framework region by second site changes can occur, for example, by imparting or 5 translating conformational changes from the second site changes to the CDR binding pocket or to the framework regions. The second site changes can occur in any of the framework regions, including for example, framework regions 1 through 4 as well as in any of the three CDR 10 regions. An advantage of the methods of the invention is that no prior information is required to assess which amino acid positions or changes can be inherently beneficial or detrimental, or which positions or changes can be further augmented by second site changes. 15 Instead, by selecting relevant amino acid positions or subsets thereof in the acceptor variable region framework and CDRs, and generating a diverse population containing amino acid variants at these positions, combinations of beneficial changes occurring at the selected positions 20 will be identified by screening for increased or optimized binding affinity of the CDR graft variable region. Such beneficial combinations will include the unveiling of inherently beneficial residues, neutralization of inherently detrimental residues and the 25 augmentation of parent residues or functionally neutral changes.

Following selection of relevant amino acid positions in the framework regions and in the donor CDRs as described previously, amino acid changes at some or all of the selected positions are incorporated into encoding nucleic acids for the acceptor variable region framework and donor CDRs, respectively. Simultaneous with incorporating the encoding amino acid changes at the selected positions, the encoding nucleic acids sequences

29

for each of the donor CDRs, including selected changes, are also incorporated into the acceptor variable region framework encoding nucleic acid to generate a population of altered variable region encoding nucleic acids.

5 An altered variable region of the invention will contain at least one framework position which variably incorporates different amino acid residues and at least one CDR position which variably incorporates different amino acid residues as described previously. 10 The variability at any or all of the altered positions can range from a few to a plurality of different amino acid residues, including all twenty naturally occurring amino acids or functional equivalents and analogues thereof. The different species of the altered variable 15 region containing the variable amino acid residues at one or more positions within the framework and CDR regions will make up the population for which to screen for an altered variable region having binding affinity substantially the same or greater than the donor CDR 20 variable region.

Selection of the number and location of the amino acid positions to vary is flexible and can depend on the intended use and desired efficiency for identification of the altered variable region having substantially the same or greater binding affinity compared to the donor variable region. In this regard, the greater the number of changes that are incorporated into a altered variable region population, the more efficient it is to identify at least one species that exhibits substantially the same or greater binding affinity as the donor. Alternatively, where the user has empirical or actual data to the affect that certain amino acid residues or positions contribute disproportionally

30

to binding affinity, then it can be desirable to produce a limited population of altered variable regions which focuses on changes within or around those identified residues or positions.

For example, if CDR grafted variable regions are desired, a large, diverse population of altered variable regions can include all the non-identical framework region positions between the donor and acceptor framework and all single CDR amino acid position changes. 10 Alternatively, a population of intermediate diversity can include subsets, for example, of only the proximal nonidentical framework positions to be incorporated together with all single CDR amino acid position changes. diversity of the above populations can be further 15 increased by, for example, additionally including all pairwise CDR amino acid position changes. In contrast, populations focusing on predetermined residues or positions which incorporate variant residues at as few as one framework and one CDR amino acid position can 20 similarly be constructed for screening and identification of an altered antibody variable region of the invention. As with the above populations, the diversity of such focused populations can be further increased by additionally expanding on the positions selected for 25 change to include other relevant positions in either or both of the framework and CDR regions. There are numerous other combinations ranging from few changes to many changes in either or both of the framework regions and CDRs that can additionally be employed, all of which 30 will result in a population of altered variable regions that can be screened for the identification of at least one CDR grafted altered variable region of the invention. Those skilled in the art will know, or can determine, which selected residue positions in the framework or

31

donor CDRs, or subsets thereof, can be varied to produce a population for screening and identification of a altered antibody of the invention given the teachings and quidance provided herein.

Simultaneous incorporation of all of the CDR encoding nucleic acids and all of the selected amino acid position changes can be accomplished by a variety of methods known to those skilled in the art, including for example, recombinant and chemical synthesis. For example, simultaneous incorporation can be accomplished by, for example, chemically synthesizing the nucleotide sequence for the acceptor variable region, fused together with the donor CDR encoding nucleic acids, and incorporating at the positions selected for harboring variable amino acid residues a plurality of corresponding amino acid codons.

One such method well known in the art for rapidly and efficiently producing a large number of alterations in a known amino acid sequence or for 20 generating a diverse population of variable or random sequences is known as codon-based synthesis or mutagenesis. This method is the subject matter of U.S. Patent Nos. 5,264,563 and 5,523,388 and is also described in Glaser et al. J. Immunology 149:3903 (1992). Briefly, 25 coupling reactions for the randomization of, for example, all twenty codons which specify the amino acids of the genetic code are performed in separate reaction vessels and randomization for a particular codon position occurs by mixing the products of each of the reaction vessels. 30 Following mixing, the randomized reaction products corresponding to codons encoding an equal mixture of all twenty amino acids are then divided into separate reaction vessels for the synthesis of each randomized

32

codon at the next position. For the synthesis of equal frequencies of all twenty amino acids, up to two codons can be synthesized in each reaction vessel.

Variations to these synthesis methods also 5 exist and include for example, the synthesis of predetermined codons at desired positions and the biased synthesis of a predetermined sequence at one or more codon positions. Biased synthesis involves the use of two reaction vessels where the predetermined or parent 10 codon is synthesized in one vessel and the random codon sequence is synthesized in the second vessel. The second vessel can be divided into multiple reaction vessels such as that described above for the synthesis of codons specifying totally random amino acids at a particular 15 position. Alternatively, a population of degenerate codons can be synthesized in the second reaction vessel such as through the coupling of NNG/T nucleotides where N is a mixture of all four nucleotides. Following synthesis of the predetermined and random codons, the 20 reaction products in each of the two reaction vessels are mixed and then redivided into an additional two vessels for synthesis at the next codon position.

A modification to the above-described codon-based synthesis for producing a diverse number of variant sequences can similarly be employed for the production of the variant populations described herein. This modification is based on the two vessel method described above which biases synthesis toward the parent sequence and allows the user to separate the variants into populations containing a specified number of codon positions that have random codon changes.

. 33

Briefly, this synthesis is performed by continuing to divide the reaction vessels after the synthesis of each codon position into two new vessels. After the division, the reaction products from each 5 consecutive pair of reaction vessels, starting with the second vessel, is mixed. This mixing brings together the reaction products having the same number of codon positions with random changes. Synthesis proceeds by then dividing the products of the first and last vessel 10 and the newly mixed products from each consecutive pair of reaction vessels and redividing into two new vessels. In one of the new vessels, the parent codon is synthesized and in the second vessel, the random codon is synthesized. For example, synthesis at the first codon 15 position entails synthesis of the parent codon in one reaction vessel and synthesis of a random codon in the second reaction vessel. For synthesis at the second codon position, each of the first two reaction vessels is divided into two vessels yielding two pairs of vessels. 20 For each pair, a parent codon is synthesized in one of the vessels and a random codon is synthesized in the second vessel. When arranged linearly, the reaction products in the second and third vessels are mixed to bring together those products having random codon 25 sequences at single codon positions. This mixing also reduces the product populations to three, which are the starting populations for the next round of synthesis. Similarly, for the third, fourth and each remaining position, each reaction product population for the 30 preceding position are divided and a parent and random codon synthesized.

Following the above modification of codon-based synthesis, populations containing random codon changes at one, two, three and four positions as well as others can

34

be conveniently separated out and used based on the need of the individual. Moreover, this synthesis scheme also allows enrichment of the populations for the randomized sequences over the parent sequence since the vessel containing only the parent sequence synthesis is similarly separated out from the random codon synthesis.

Other methods well known in the art for producing a large number of alterations in a known amino acid sequence or for generating a diverse population of variable or random sequences include, for example, degenerate or partially degenerate oligonucleotide synthesis. Codons specifying equal mixtures of all four nucleotide monomers, represented as NNN, results in degenerate synthesis. Whereas partially degenerate synthesis can be accomplished using, for example, the NNG/T codon described previously. Other method well know in the art can alternatively be used such as the use of statistically predetermined, or varigated, codon synthesis which is the subject matter of U.S. Patent Nos. 5,223,409 and 5,403,484.

Once the populations of altered variable region encoding nucleic acids have been constructed as described above, they can be expressed to generate a population of altered variable region polypeptides that can be screened for binding affinity. For example, the altered variable region encoding nucleic acids can be cloned into an appropriate vector for propagation, manipulation and expression. Such vectors are known or can be constructed by those skilled in the art and should contain all expression elements sufficient for the transcription, translation, regulation, and if desired, sorting and secretion of the altered variable region polypeptides. The vectors also can be for use in either procaryotic or

WO 01/27160

PCT/US00/28435

eukaryotic host systems so long as the expression and regulatory elements are of compatible origin. The expression vectors can additionally included regulatory elements for inducible or cell type-specific expression.

5 One skilled in the art will know which host systems are compatible with a particular vector and which regulatory or functional elements are sufficient to achieve expression of the polypeptides in soluble, secreted or cell surface forms.

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10 Appropriate host cells, include for example, bacteria and corresponding bacteriophage expression systems, yeast, avian, insect and mammalian cells. Methods for recombinant expression, screening and purification of populations of altered variable regions 15 or altered variable region polypeptides within such populations in various host systems are well known in the art and are described, for example, in Sambrook et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, New York (1992) and in Ansubel et al., 20 Current Protocols in Molecular Biology, John Wiley and Sons, Baltimore, MD (1998). The choice of a particular vector and host system for expression and screening of altered variable regions will be known by those skilled in the art and will depend on the preference of the user. 25 A specific example of the expression of recombinant altered variable region polypeptides is additionally described below in the Examples. Moreover, expression of diverse populations of hetereomeric receptors in either soluble or cell surface form using filamentous 30 bacteriophage vector/host systems is well known in the art and is the subject matter of U.S. Patent No. 5,871,974.

36

The expressed population of altered variable region polypeptides can be screened for the identification of one or more altered variable region species exhibiting binding affinity substantially the 5 same or greater than the donor CDR variable region. Screening can be accomplished using various methods well known in the art for determining the binding affinity of a polypeptide or compound. Additionaly, methods based on determining the relative affinity of binding molecules to 10 their partner by comparing the amount of binding between the altered variable region polypeptides and the donor CDR variable region can similarly be used for the identification of species exhibiting binding affinity substantially the same or greater than the donor CDR 15 variable region. All of such methods can be performed, for example, in solution or in solid phase. Moreover, various formats of binding assays are well known in the art and include, for example, immobilization to filters such as nylon or nitrocellulose; two-dimensional arrays, 20 enzyme linked immunosorbant assay (ELISA), radioimmune assay (RIA), panning and plasmon resonance. Such methods can be found described in, for example, Sambrook et al., supra, and Ansubel et al.

For the screening of populations of

25 polypeptides such as the altered variable region
populations produced by the methods of the invention,
immobilization of the populations of altered variable
regions to filters or other solid substrate is
particularly advantageous because large numbers of

30 different species can be efficiently screened for antigen
binding. Such filter lifts will allow for the
identification of altered variable regions that exhibit
substantially the same or greater binding affinity
compared to the donor CDR variable region. Alternatively,

37

if the populations of altered variable regions are expressed on the surface of a cell or bacteriophage, for example, panning on immobilized antigen can be used to efficiently screen for the relative binding affinity of species within the population and for those which exhibit substantially the same or greater binding affinity than the donor CDR variable region.

Another affinity method for screening populations of altered variable regions polypeptides is a 10 capture lift assay that is useful for identifying a binding molecule having selective affinity for a ligand (Watkins et. al., (1997)). This method employs the selective immobilization of altered variable regions to a solid support and then screening of the selectively 15 immobilized altered variable regions for selective binding interactions against the cognate antigen or binding partner. Selective immobilization functions to increase the sensitivity of the binding interaction being measured since initial immobilization of a population of 20 altered variable regions onto a solid support reduces non-specific binding interactions with irrelevant molecules or contaminants which can be present in the reaction.

Another method for screening populations or for measuring the affinity of individual altered variable region polypeptides is through surface plasmon resonance (SPR). This method is based on the phenomenon which occurs when surface plasmon waves are excited at a metal/liquid interface. Light is directed at, and reflected from, the side of the surface not in contact with sample, and SPR causes a reduction in the reflected light intensity at a specific combination of angle and wavelength. Biomolecular binding events cause changes in

WO 01/27160

38

PCT/US00/28435

the refractive index at the surface layer, which are detected as changes in the SPR signal. The binding event can be either binding association or disassociation between a receptor-ligand pair. The changes in refractive index can be measured essentially instantaneously and therefore allows for determination of the individual components of an affinity constant. More specifically, the method enables accurate measurements of association rates  $(k_{on})$  and disassociation rates  $(k_{off})$ .

Measurements of kon and koff values can be 10 advantageous because they can identify altered variable regions or optimized variable regions that are therapeutically more efficacious. For example, an altered variable region, or heteromeric binding fragment 15 thereof, can be more efficacious because it has, for example, a higher kon valued compared to variable regions and heteromeric binding fragments that exhibit similar binding affinity. Increased efficacy is conferred because molecules with higher kon values can specifically 20 bind and inhibit their target at a faster rate. Similarly, a molecule of the invention can be more efficacious because it exhibits a lower koff value compared to molecules having similar binding affinity. Increased efficacy observed with molecules having lower 25 koff rates can be observed because, once bound, the molecules are slower to dissociate from their target. Although described with reference to the altered variable regions and optimized variable regions of the invention including, heteromeric variable region binding fragments 30 thereof, the methods described above for measuring associating and disassociation rates are applicable to essentially any antibody or fragment thereof for identifying more effective binders for therapeutic or diagnostic purposes.

39

Methods for measuring the affinity, including association and disassociation rates using surface plasmon resonance are well known in the arts and can be found described in, for example, Jönsson and Malmquist,

5 Advances in Biosnsors, 2:291-336 (1992) and Wu et al.

Proc. Natl. Acad. Sci. USA, 95:6037-6042 (1998).

Moreover, one apparatus well known in the art for measuring binding interactions is a BIAcore 2000 instrument which is commercially available through

10 Pharmacia Biosensor, (Uppsala, Sweden).

Using any of the above described screening methods, as well as others well known in the art, an altered variable region having binding affinity substantially the same or greater than the donor CDR 15 variable region is identified by detecting the binding of at least one altered variable region within the population to its antigen or cognate ligand. Additionally, the above methods can alternatively be modified by, for example, the addition of substrate and 20 reactants, to identify using the methods of the invention, altered variable regions having catalytic activity substantially the same or greater that the donor CDR variable region within the populations. Comparision, either independently or simultaneously in the same 25 screen, with the donor variable region will identify those binders that have substantially the same or greater binding affinity as the donor. Those skilled in the art will know, or can determine using the donor variable region, binding conditions which are sufficient to 30 identify selective interactions over non-specific binding.

Detection methods for identification of binding species within the population of altered variable regions

40

can be direct or indirect and can include, for example, the measurement of light emission, radioisotopes, colorimetric dyes and fluorochromes. Direct detection includes methods that operate without intermediates or 5 secondary measuring procedures to assess the amount of bound antigen or ligand. Such methods generally employ ligands that are themselves labeled by, for example, radioactive, light emitting or fluorescent moieties. In contrast, indirect detection includes methods that 10 operate through an intermediate or secondary measuring procedure. These methods generally employ molecules that specifically react with the antigen or ligand and can themselves be directly labeled or detected by a secondary reagent. For example, a antibody specific for a ligand 15 can be detected using a secondary antibody capable of interacting with the first antibody specific for the ligand, again using the detection methods described above for direct detection. Indirect methods can additionally employ detection by enzymatic labels. Moreover, for the 20 specific example of screening for catalytic antibodies, the disappearance of a substrate or the appearance of a product can be used as an indirect measure of binding affinity or catalytic activity.

Isolated variable regions exhibit binding
25 affinity as single chains, in the absence of assembly into a heteromeric structure with their respective V<sub>H</sub> or V<sub>L</sub> subunits. As such, populations of V<sub>H</sub> and V<sub>L</sub> altered variable regions polypeptides can be expressed alone and screened for binding affinity having substantially the
30 same or greater binding affinity compared to the CDR donor V<sub>H</sub> or V<sub>L</sub> variable region. Alternatively, populations of V<sub>H</sub> and V<sub>L</sub> altered variable regions polypeptides can be coexpressed so that they selfassemble into heteromeric altered variable region binding

41

fragments. The heteromeric binding fragment population can then be screened for species exhibiting binding affinity substantially the same or greater than the CDR donor variable region binding fragment. A specific example of the coexpression and self-assembly of populations  $V_R$  and  $V_L$  altered variable regions into hetermeric populations is described further below in the Examples.

Therefore, the invention provides a method of 10 simultaneously grafting and optimizing the binding affinity of a variable region binding fragment. method consists of: (a) constructing a population of altered heavy chain variable region encoding nucleic acids consisting of an acceptor variable region 15 framework, containing donor CDRs and a plurality of different amino acids at one or more framework region and CDR amino acid positions; (b) coexpressing the populations of heavy and light chain variable region encoding nucleic acids to produce diverse combinations of 20 heteromeric variable region binding fragments, and (c) identifying one or more heteromeric variable region binding fragments having affinity substantially the same or greater than the donor CDR heteromeric variable region binding fragment.

The invention additionally provides a method of optimizing the binding affinity of an antibody variable region. The consists of: (a) constructing a population of antibody variable region encoding nucleic acids, said population comprising two or more CDRs containing a plurality of different amino acids at one or more CDR amino acid positions; (b) expressing said population of variable region encoding nucleic acids, and (c) identifying one or more variable regions having binding

42

affinity substantially the same or greater than the donor CDR variable region.

The methods described above, for conferring donor CDR binding affinity onto an antibody acceptor 5 variable region framework and for simultaneously grafting and optimizing the binding affinity of a heteromeric variable region binding fragment, can additionally be employed to modify or optimize the binding affinity of a variable region or a heteromeric variable region binding 10 fragment. Similar to the previously described methods, the method for modifying or optimizing binding affinity involves the selection of relevant amino acid positions and the construction, expression and screening of variable region populations containing variable amino 15 acid residues at all or a fraction of the selected positions. However, for optimization of binding affinity it is not necessary to vary amino acid positions in the framework regions. Instead, all that is required is to alter one or more amino acid positions in two or more CDR 20 regions. Changing the CDR amino acid residues directly effects the binding affinity. Once a population containing variable amino acid residues incorporated in two or more CDRs is produced, all that is necessary is to screen the population for species that contain the 25 desired binding affinity modification. All of the criteria for selecting relevant amino acid positions described previously are applicable for use in this mode of the method. Therefore, the methods for modifying or optimizing the binding affinity of a variable region or a 30 heteromeric variable region binding fragment by altering one or more amino acid positions in two or more CDR regions are applicable to essentially any variable region, grafted variable region as well as applicable to

43

the altered and optimized variable regions of the invention.

Moreover, by incorporating variable amino acid residues in two or more CDRs when employing the methods conferring donor CDR binding affinity onto an acceptor framework, this method of modifying binding affinity is therefore useful for simultaneously optimizing the binding affinity of a grafted antibody. Employing the methods for simultaneously grafting and optimizing, or for optimizing, it is possible to generate heteromeric variable region binding fragments having increases in affinities of greater than 5-, 8- and 10-fold. In particular, heteromeric variable region binding fragments can be generated having increases in affinities of greater than 12-, 15- 20- and 25-fold as well as affinities greater than 50-, 100- and 1000-fold compared to the donor or parent molecule.

Additionally, the methods described herein for optimizing are also are applicable for producing

20 catalytic heteromeric variable region fragments or for optimizing their catalytic activity. Catalytic activity can be optimized by changing, for example, the on or off rate, the substrate binding affinity, the transition state binding affinity, the turnover rate (kcat) or the

25 Km. Methods for measuring these characteristics are well known in the art. Such methods can be employed in the screening steps of the methods described above when used for optimizing the catalytic activity of a heteromeric variable region binding fragment.

30 The methods for conferring donor CDR binding affinity onto an antibody acceptor variable region framework described previously are applicable for use

with essentially any distinguishable donor and acceptor pair. Many applications of the methods will be for the production and optimization of variable region binding fragments having human acceptor frameworks due to the therapeutic importance of such molecules in the treatment of human diseases. However, the method are applicable for conferring donor CDR binding affinity onto an acceptor originating from the same or a divergent species as the CDR donor variable region so long as the framework regions between the donor and acceptor variable regions are distinct. Therefore, the invention included altered variable regions having acceptor frameworks derived from human, mouse, rat, rabbit, goat and chicken, for example.

44

Additionally, the methods for conferring donor 15 CDR binding affinity onto an antibody acceptor variable region framework are applicable for grafting CDRs as described by Kabat et al., supra, Chothia et al., supra or MacCallum et al., supra. The methods similarly can be used for grafting into an acceptor framework overlapping 20 regions or combinations of CDR as described by these authors. Generally, the methods will graft variable region CDRs by identifying the boundries described by one of the CDR definitions known in the art and set forth herein. However, because the methods are directed to 25 constructing and screening populations of CDR grafted altered variable regions which incorporate relevant amino acid position changes in both the framework and CDR regions, and such variations can, for example, compensate or augment amino acid changes elsewhere in the variable 30 region, the exact boundry of a particular CDR or set of variable region CDRs can be varied. Therefore, the exact CDR region to graft, whether it is the region described by Kabat et al., Chothia et al. or MacCallum et al., or

45

any combination thereof, will essentially depend on the preference of the user.

Similarly, the methods described previously for optimizing the binding affinity of an antibody also are 5 applicable for use with essentially any variable region for which an encoding nucleic acid is, or can be made available. As with the methods for conferring donor CDR binding affinity, many applications of the methods for optimizing binding affinity will be for modifying the 10 binding affinity of CDR grafted variable regions having human frameworks. Again, such molecules are significantly less antigenic in human patients and therefore therapeutically valuable in the treatment of human diseases. However, the methods of the invention 15 for optimizing the binding affinity of a variable region are applicable to all species of variable regions. Therefore, the invention includes binding affiity optimization of variable regions derived from human, mouse, rat, rabbit, goat and chicken, for example.

The methods of the invention have been described with reference to variable regions and heteromeric variable region binding fragments. Given these descriptions and teaching herein, those skilled in the art will understand that all of such methods are applicable to whole antibodies and functional fragments thereof as well as to regions and functional domains other than the antigen binding variable region of antibodies. Moreover, the methods described herein are further applicable to molecules other than antibodies, variable regions and other antibody functional domains. Given the teachings of the invention, those skilled in the art will know how to apply the methods of simultaneously constructing hybrid molecules and

WO 01/27160

46

PCT/US00/28435

maintaining or optimizing the binding affinity or catalytic activity of a target molecule, as well as how to apply the methods of optimizing the binding affinity or catalytic activity to a variety of different types and 5 classes of polypeptides and proteins.

The methods for optimizing the binding affinity of an antibody variable region can include the selection of relevant acceptor framework and donor CDR amino acid 10 positions to be altered. Amino acid residues selected for alteration during binding affinity optimization are typically amino positions predicted to be relatively important for structure or function. Criteria that can be used for identifying amino positions to be altered 15 include, for example, conservation of amino acids among polypeptide subfamily members and knowledge that particular amino acids are predicted to be important in polypeptide conformation or structure, as described above. Alternatively, potentially important framework 20 residues that differ between acceptor framework and donor CDR can be characterized without structural information by synthesizing and expressing a combinatorial antibody library that contains all possible combinations of amino acids in framework positions to be optimized.

The invention provides a method for identifying one or more functional amino acid positions of a polypeptide. The method consists of (a) constructing a population of nucleic acids encoding a population of altered polypeptides containing substitutions of one or more amino acid positions within a polypeptide;

(b) expressing the population of nucleic acids;

(c) identifying nucleic acids encoding altered polypeptides having a functional activity of the

polypeptide; (d) sequencing a subset of nucleic acids

47

encoding altered polypeptides having a functional activity, and (e) comparing an amino acid position in a polypeptide corresponding to an amino acid position in the subset of altered polypeptides wherein an amino acid position exhibiting a biased representation of amino acid residues indicates a functional amino acid position in the polypeptide.

The method of the invention directed to identifying a functional amino acid position in a polypeptide involves substituting one or more amino acid positions in a polypeptide with a plurality of amino acid residues, as described previously for optimizing the binding affinity of an antibody, and identifying altered polypeptides having an activity that is substantially the same or greater than the parent polypeptide. Functional amino acid positions identified using the methods of the invention are amino acid positions important for a conformation, functional activity or structure of a polypeptide. Functional activities of a polypeptide can include, for example, binding affinity to a substrate, ligand, or other interacting molecule, and catalytic activity.

The identification of functional amino acid positions in a polypeptide involves constructing a population of nucleic acids encoding a population of altered polypeptides containing amino acid substitutions at specific amino acid positions. Substituted amino acids include all twenty naturally occurring amino acid residues or a subset of amino acid residues, as described previously in detail. Nucleic acid populations can be constructed by any method known in the art and as described previously. A population of nucleic acids encoding altered polypeptides is expressed in an

48

appropriate host cell, and a functional activity of altered polypeptides is detected and compared with that of the polypeptide. Any method known in the art that is appropriate for determining a polypeptide functional activity can be used to compare polypeptide and altered polypeptide functional activities.

A subset of nucleic acids encoding altered polypeptides having a functional activity that is substantially the same or greater than that of the 10 polypeptide is sequenced. A subset can include a few molecules to many members constituting the population of nucleic acids encoding altered polypeptides. For example, a subset can consist of about 2-5, 6-10, 10-20, and 21 or greater members of the population. The actual 15 number sequenced will vary with the total size of the nucleic acid population. Generally, however, a subset of about 15-25 and typically about 20 members is sufficient in order to identify functional amino acids.

Amino acid residues at substituted positions in
the polypeptide are compared to the corresponding
position in altered polypeptides. An amino acid position
that contains the same amino acid or a conservative
substitution among the population of altered polypeptides
exhibits biased representation of that amino acid
residue. Biased representation indicates that a
particular amino acid is required for polypeptide
function. Amino acid positions that are biased are
therefore considered important for functional activity of
a polypeptide. Amino acid positions that contain a
variety of substituted amino acids are unbiased and
considered not important or less important for a
polypeptide function.

49

The method of identifying an amino acid position important for polypeptide function is useful for a variety of applications, such as, for example, the determination of a consensus sequence of amino acids

5 important for a polypeptide functional activity. A consensus sequence is useful for the optimization of a polypeptide function because amino acid positions determined to be important for functional activity can be unaltered while amino acid positions not important for activity can be varied. Polypeptide functions that can be optimized using the method of the invention include, for example, catalytic activity, polypeptide conformation and binding affinity.

The identification of a functional amino acid

position in a polypeptide can be applied to determining a
consensus sequence of amino acids that impart a
particular activity to a polypeptide. For example, a
consensus sequence that provides a catalytic activity to
an enzyme can be determined using the methods of the

invention. To identify amino acid positions that are
important or critical to catalytic activity of an enzyme,
one or more of amino acid positions are substituted with
a plurality of amino acid substitutions, as described
previously. A nucleic acid population encoding altered
enzyme polypeptides is constructed and expressed in host
cells. The catalytic activity of altered enzymes is
measured and compared with a parent enzyme or other
catalytically active form of the enzyme.

Nucleic acids encoding a subset of altered

30 enzyme polypeptides identified by functional activity are sequenced, and the amino acid sequences of altered polypeptides are compared. Amino acid positions that contain a particular amino acid or a conservative

50

substitution are determined to be important for a catalytic activity of the enzyme. A sequence of amino acids determined to be biased in a polypeptide can thus provide a consensus sequence that defines amino acid 5 positions required for catalytic activity. A consensus sequence of residues important for various aspects of catalytic activity such as, for example, substrate binding, proper active site conformation, and co-factor binding can be identified using the methods of the invention by measuring enzyme catalytic activity, as described above.

Similarly, a consensus sequence associated with a particular conformation of a polypeptide can be determined using the method of the invention in

15 essentially the same manner as described above for polypeptide catalytic activity. The amino acid positions that have functional roles in a polypeptide conformation can be determined so long as a particular conformation state can be detected and compared between a polypeptide and an altered polypeptide. For example, a consensus sequence of a polypeptide conformation that confers a particular functional activity to a polypeptide or a particular structural feature to a polypeptide can be determined using the methods of the invention. A

25 structural feature can include, for example, the exposure of a certain amino acid on the surface of a polypeptide.

A consensus sequence of amino acid positions in a polypeptide important for binding affinity can also be determined using the methods of the invention. The binding affinities of polypeptides include, for example, the binding affinity between two or more polypeptides in a protein-protein interaction and the binding affinity between a polypeptide and a substrate. For example, a

51

consensus sequence for the binding affinity of an antibody for an antigen can be determined, and can be applied to the process of antibody humanization.

The identification of a functional amino acid 5 position in a polypeptide can be applied to determining the consensus sequence for a humanized version of an antibody that preserves the binding activity of the parent antibody. For example, a library containing all possible combinations of human template and murine parent 10 antibody residues in a selected number of amino positions can be synthesized by method known in the art, for example, using codon-based mutagenesis as described previously. Framework polypeptides containing amino acid substitutions can then be screened by functional binding 15 to identify altered framework polypeptides that have a binding affinity substantially the same as the parent antibody. Of the amino acid positions altered, only a small percentage of framework positions are typically critical for antibody binding activity. Therefore, a low 20 throughput screening method of identifying active humanized framework variants can be used. Sequencing of nucleic acids encoding humanized frameworks displaying a functional activity of the parent antibody is then used to identify altered polypeptides having significant bias 25 toward murine human residues. Thus, a consensus humanization sequence for maintaining full binding activity of an antibody can be prepared by using murine CDRs grafted onto a human template on which amino acid positions are changed to the corresponding residue 30 determined to be important for binding activity.

It is understood that modifications which do not substantially affect the activity of the various embodiments of this invention are also included within

52

the definition of the invention provided herein. Accordingly, the following examples are intended to illustrate but not limit the present invention.

#### EXAMPLE I

# 5 Simultaneous Humanization and Affinity Maturation of an Anti-CD40 Antibody

This example shows the simultaneous humanization and affinity maturation of the murine mAb 40.2.220, directed against the CD40 receptor.

10 The CD40 receptor is a potential therapeutic target for several diseases. For example, the interaction of the CD40 receptor and its ligand, gp39, serves a critical role in both humoral and cell-mediated immune responses (Foy et. al., (1996)). Immunological 15 rejection of organs from genetically non-identical individuals, termed graft-versus-host-disease (GVHD), is mediated through T cell-dependent mechanisms. In vivo administration of an anti-gp39 mAb blocks GVHD in mice and inhibits many of the GVHD-associated phenomena (Durie 20 et. al., 1994)), providing evidence that the CD40/gp39 interaction plays a critical role in the development of GVHD. More recently, inhibition of the CD40/gp39 interaction in vivo in hyperlipidemic mice fed a high cholesterol diet limited atherosclerosis, suggesting that 25 CD40 signalling may also play a role in atherogenesis (Mach et. al., (1998)). In addition, the CD40 receptor is overexpressed on hematologic malignancies (Uckun et. al., 1990)) and certain carcinomas (Stamenkovic et. al., (1989)) and thus, may serve as a target for cytotoxic 30 agents. An anti-CD40 single chain antibody-toxin fusion was cytotoxic against CD40-expressing malignant cells in vitro (Francisco et. al., (1995)) and was efficacious in

53

treating human non-Hodgkin's lymphoma xenografted SCID mice (Francisco et. al., (1997)).

Codon-based mutagenesis (Glaser et. al., (1992)) was used to create libraries of LCDR3, HCDR3 and framework region variants of mAb 40.2.220 sequences. Libraries composed of framework region variants alone and in combination with HCDR3 variants and with HCDR3 and LCDR3 variants together were screened for high affinity variants. It was demonstrated that in combination higher affinity variants were obtained than those obtained when codon-based mutagenesis was applied independently thus showing (1) higher affinity variants that could only be obtained by the use of codon-based mutagenesis simultaneously on disparate regions of the mAb and (2) the use of codon-based mutagenesis to uncover potential direct interactions between disparate regions of a mAb.

A vector for the production of a chimeric anti-CD40 murine mAb 40.2.220 was constructed. Based on the sequence of anti-CD40 murine mAb 40.2.220 (provided by 20 Dr. D. Hollenbaugh, Bristol-Myers Squibb, Princeton, NJ) overlapping oligonucleotides encoding  $V_{\text{H}}$  and  $V_{\text{L}}$  (69-75 bases in length) were synthesized and purified. The variable H and L domains were synthesized separately by combining 25 pmol of each of the overlapping 25 oligonucleotides with Pfu DNA polymerase (Stratagene) in a 50  $\mu$ l PCR reaction consisting of 5 cycles of: denaturing at 94°C for 20 sec, annealing at 50°C for 30 sec, ramping to 72°C over 1 min, and maintaining at 72°C for 30 sec. Subsequently, the annealing temperature was 30 increased to 55°C for 25 cycles. A reverse primer and a biotinylated forward primer were used to further amplify 1  $\mu$ l of the fusion product in a 100  $\mu$ l PCR reaction using the same program. The products were purified by agarose

54

gel electrophoresis, electroeluted, and phosphorylated by T4 polynucleotide kinase (Boehringer Mannheim) and were then incubated with streptavidin magnetic beads (Boehringer Mannheim) in 5 mM Tris-Cl, pH 7.5, 0.5 mM 5 EDTA, 1 M NaCl, and 0.05% Tween 20 for 15 min at 25°C. The beads were washed and the non-biotinylated, minus strand DNA was eluted by incubating with 0.15 M NaOH at 25°C for 10 min. Chimeric anti-CD40 Fab was synthesized in a modified M13IX104 phage vector (Kristensson et. al., 10 1995)), termed M13IX104CS, by hybridization mutagenesis (Rosok et. al., (1996); Kunkel, (1985)) using the  $V_H$  and V, oligonucleotides in 3-fold molar excess of the uridinylated vector template. The M13IX104 vector was modified by replacing cysteine residues at the end of the 15 kappa and y1 constant regions with serine. The reaction was electroporated into DH10B cells and titered onto a lawn of XL-1 Blue.

The murine anti-CD40 mAb variable region framework sequences were used to identify the most

20 homologous human germline sequences. The H chain framework residues were 74% identical to human germline VH7 (7-4.1) and JH4 sequences while the L chain was 75% identical to the corresponding human germline VKIII (L6) and JK4 sequences. Alignment of the H and L chain

25 variable sequences is shown in Fig. 1. CDR residues, as defined by Kabat et. al. (1977, 1991), are underlined and were excluded from the homology analysis. Differences in sequence are indicated by vertical lines and framework positions characterized in the combinatorial expression

30 library are marked with an asterisk. Framework residues that differed between the murine mAb and the human templates were assessed individually.

55

Based on structural and sequence analysis, antibody CDRs with the exception of HCDR3 display a limited number of main chain conformations termed canonical structures (Chothia & Lesk, (1987); Chothia et. 5 al., (1989)). Moreover, certain residues critical for determining the main chain conformation of the CDR loops have been identified (Chothia & Lesk, (1987); Chothia et. al., (1989)). Canonical framework residues of murine anti-CD40 were identified therefore, and it was determined that amino acids at all critical canonical positions within the H and L chain frameworks of the human templates were identical to the corresponding murine residues.

Surface-exposed murine amino acids not normally 15 found in human antibodies are likely to contribute to the immunogenicity of the humanized mAb (Padlan, (1991)). Therefore, framework residues differing between murine anti-CD40 and the human templates were analyzed and based on solvent exposure were predicted to be buried or 20 located on the surface of the antibody (Padlan, (1991)). Solvent-exposed framework residues distal to the CDRs were not expected to contribute to antigen binding significantly and thus, with the exception of two H chain residues all were changed to the corresponding human 25 amino acid to decrease potential immunogenicity. H chain residues 28 and 46 were predicted to be solvent exposed. However, H28 is located within the HCDR1 region as defined by Chothia & Lesk (1987) and potentially interacts with the antigen. In addition, the lysine at 30 H46 in the murine mAb is somewhat unusual and . significantly different from the glutamic acid of the human template. Therefore, the murine and human residues at H28 and H46 were expressed in the combinatorial library (Fig. 1, asterisks).

56

The remaining differing framework residues, all predicted to be mostly buried within the antibody, were evaluated for: (1) proximity to CDRs, (2) potential to contact the opposite domain in the  $V_R-V_H$  interface, (3) 5 relatedness of the differing amino acids, and (4) predicted importance in modulating CDR activity as defined by Studnicka et. al., (1994). The majority of L chain framework differences in buried residues were related amino acids at positions considered not likely to 10 be directly involved in the conformation of the CDR. However, L49 is located adjacent to LCDR2, potentially contacts the V<sub>B</sub> domain, is unrelated to the human residue, and may be involved in determining the conformation of LCDR2. For these reasons, the murine and human amino 15 acids at L49 were both expressed in the combinatorial framework library (Fig. 1, asterisk).

Analysis of the murine H chain sequence and the human template was performed. Residue H9 is a proline in the murine mAb while the human template contains an 20 unrelated serine residue. Position H9 can also play a role in modulating the conformation of the CDR and thus, was selected as a combinatorial library site (Fig. 1, asterisks). The remaining buried framework residues that differed between murine anti-CD40 and the H chain 25 template were at framework positions 38, 39, 48, and 91. Murine anti-CD40 mAb contained glutamine and glutamic acid at H38 and H39, respectively, while the human template contained arginine and glutamine. Residue H38 is in proximity to the HCDR1, the glutamine→arginine 30 change is non-conserved, and expression of glutamine at this site in murine Abs is somewhat unusual. Similarly, glutamic acid→glutamine is a non-conservative difference for buried amino acids, H39 is a potential  $V_{\kappa}$  contact residue, and glutamic acid is somewhat unusual in murine

57

mAbs. Residue H48 is in close proximity to HCDR2 and H91 is predicted to be a high risk site (Studnicka et. al., (1994); Harris & Bajorath, (1995)) that potentially contacts the  $V_K$  domain. Thus, both murine and human residues were expressed at H38, 39, 48, and 91 (Fig. 1, asterisks).

The combinatorial framework library (Hu I) was synthesized by the same method used to construct the chimeric anti-CD40, with modifications. Overlapping oligonucleotides encoding the framework regions of the H and L variable domains of the human template and the murine anti-CD40 CDRs as defined by Kabat et. al. (1977, 1991) were synthesized. Among these, degenerate oligonucleotides encoding both the murine and the human amino acids at seven V<sub>H</sub> and one V<sub>K</sub> framework position as selected above were synthesized (Fig. 1 residues marked with asterisk). All of these sites were characterized by synthesizing a combinatorial library that expressed all possible combinations of the murine and human amino acids found at these residues. The total diversity of this library, termed Hu I, was 28 or 256 variants (Table I).

The Hu I combinatorial library was first screened by an ELISA that permits the rapid assessment of the relative affinities of the variants (Watkins et. al., (1997)). Briefly, microtiter plates were coated with 5 µg/ml goat anti-human kappa (Southern Biotechnology) and blocked with 3% BSA in PBS. Certain Fabs were cloned into an expression vector under the control of the arabinose-regulated BAD promoter. In addition, a six-histidine tag was fused to the carboxyl-terminus of the H chain to permit purification with nickel-chelating resins. Purified Fab was quantitated as described (Watkins et. al., 1997). Next, 50 µl Fab from the

58

Escherichia coli culture supernatant or from the cell lysate, was incubated with the plate 1 h at 25°C, the plate was washed three times with PBS containing 0.1% Tween 20, and incubated with 0.1 μg/ml CD40-Ig in PBS containing 1% BSA for 2 h at 25°C. The plate was washed three times with PBS containing 0.1% Tween 20 and goat anti-mouse IgG<sub>2b</sub>-alkaline phosphatase conjugate diluted 3000-fold in PBS containing 1% BSA was added for 1 h at 25°C. The plate was washed three times with PBS containing 0.1% Tween 20 and was developed as described (Watkins et. al., (1997)).

Although variants that bind the target antigen with affinities comparable to, or better than the chimeric Fab were identified, the majority of Hu I clones screened were less active than the chimeric anti-CD40 Fab. Approximately 6% of randomly selected Hu I variants displayed binding activities comparable to the chimeric Fab (data not shown). The identification of multiple Hu I variants with activity comparable to the chimeric CD40 is consistent with the interpretation that the most critical framework residues were included in the combinatorial library.

The kinetic constants for the interaction between CD40 and the anti-CD40 variants were determined by surface plasmon resonance (BIAcore). CD40-Ig fusion protein was immobilized to a (1-ethyl-3-[3-dimethylaminopropyl]-carbodiimide hydrochloride) and N-hydroxysuccinimide-activated sensor chip CM5 by injecting 8  $\mu$ l of 10  $\mu$ g/ml CD40-Ig in 10 mM sodium acetate, pH 4. CD40-Ig was immobilized at a low density (~150 RU) to prevent rebinding of Fabs during the dissociation phase. To obtain association rate constants ( $k_{\rm en}$ ), the binding rate at six different Fab

WO 01/27160

59

PCT/US00/28435

concentrations ranging from 25-600 nM in PBS was determined at a flow rate of 20  $\mu$ l/min. Dissociation rate constants ( $k_{\rm off}$ ) were the average of six measurements obtained by analyzing the dissociation phase.

5 Sensorgrams were analyzed with the BIAevaluation 3.0 program.  $K_d$  was calculated from  $K_d = k_{off}/k_{on}$ . Residual Fab was removed after each measurement by prolonged dissociation.

Figure. 2A shows bacterially-expressed chimeric 10 anti-CD40 Fab and certain variants from each of the libraries were titrated on immobilized antigen. Chimeric (filled circles), Hu I-19C11 (open circles), Hu II-CW43 (open squares), Hu III-2B8 (filled triangles), and an irrelevant (filled squares) Fab were released from the 15 periplasmic space of 15 ml bacterial cultures and serial dilutions were incubated with CD40-Ig antigen immobilized on microtiter plates. See below for description of HuII and HuIII libraries. Antibody binding was quantitated as described above. These measurements confirm the 20 identification of multiple variants with enhanced affinity. For example, clone 19C11 binds the CD40 receptor with higher affinity than the chimeric Fab, as demonstrated by the shift in the titration profile (compare open circles with filled circles). DNA 25 sequencing of 34 of the most active clones led to the identification of 24 unique framework combinations, each containing 2-6 murine framework residues (data not shown).

LCDR3 and HCDR3 contact antigen directly,

30 interact with the other CDRs, and often affect the
affinity and specificity of antibodies significantly
(Wilson & Stanfield, (1993); Padlan, (1994)). In
addition, the conformation of LCDR3 and HCDR3 are

WO 01/27160

60

PCT/US00/28435

determined in part by certain framework residues.

Therefore, to identify the most active antibody it could be beneficial to construct combinatorial libraries that optimize the third CDR of the H and L chains in conjunction with selecting the most active framework.

Codon-based mutagenesis (Glaser et. al., (1992)) was used to synthesize oligonucleotides that introduce mutations at every position in HCDR3, one at a time, resulting in the expression of all 20 amino acids at each CDR residue from Ser<sup>95</sup>-Tyr<sup>102</sup> (Fig. 1, underlined). Briefly, for library construction, the overlapping oligonucleotides encoding the framework library and non-library murine CDR were combined with 25 pmol of the oligonucleotides encoding mutated HCDR3. The pool of oligonucleotides encoding the HCDR3 library was mixed with the overlapping oligonucleotides encoding the combinatorial framework and other CDRs to generate a framework/HCDR3 library. The diversity of this library, termed Hu II, was 1.1 x 10<sup>5</sup> (Table I).

The CDR residues selected for mutagenesis of LCDR3 were Gln<sup>89</sup>-Thr<sup>97</sup> (Fig. 1, underlined).

Oligonucleotides encoding LCDR3 were designed to mutate a single CDR residue in each clone as described above for HCDR3. Oligonucleotides encoding the LCDR3, HCDR3, and the combinatorial framework were used to create a framework/HCDR3/LCDR3 library, termed Hu III. The large number of framework/CDR3 combinations resulted in a library with a complexity of 3.1 x 10<sup>7</sup> (Table I).

Table I. Summary of phage-expressed anti-CD40 antibody libraries.

Library	Library Positions	Size*	Screened <sup>t</sup>	
Hu I	framework	256	2.4 x 10 <sup>3</sup>	
Hu II	framework, HCDR3	1.1 x	$2.0 \times 10^{6}$	
		105		
Hu III	framework, HCDR3, LCDR3	3.1 x	5.5 x 10 <sup>5</sup>	
		107		

5

\*Number of unique clones based on DNA sequence.
Thirty-two codons are used to encode all 20 amino acids at each CDR position.

An additional library (Hu IV) was synthesized to further optimize the best variant (clone F4) identified from the Hu III library. Oligonucleotides encoding LCDR3, designed to mutate a single CDR residue in each clone, were synthesized by introducing NN(G/T) at each position (Glaser et. al., (1992)) and were annealed to uridinylated F4 template (Kunkel, (1985)) which already contained a 96R→W mutation in LCDR3.

Combining mutations in LCDR3 and/or HCDR3 with the framework library increased the potential diversity of humanized anti-CD40 variants from 256 to greater than 10<sup>7</sup>. In order to screen these larger libraries more efficiently a modified plaque lift assay, termed capture lift, was used (Watkins et. al., (1997)). Briefly, nitrocellulose filters (82-mm) were coated with goat anti-human kappa, blocked with 1% BSA, and were applied to an agar plate containing the phage-infected bacterial lawn. In the initial screen, phage were plated at 10<sup>5</sup> phage/100-mm plate. After the capture of phage-expressed

62

anti-CD40 variant Fabs, the filters were incubated 3 h at 25°C with 5 ng/ml CD40-Ig in PBS containing 1% BSA. The filters were rinsed four times with PBS containing 0.1% Tween 20 and were incubated with goat anti-mouse 5  $IgG_{2b}$ -alkaline phosphatase conjugate (Southern Biotechnology) diluted 3000-fold in PBS containing 1% BSA for 1 h at 25°C. The filters were washed four times with PBS containing 0.1% Tween 20 and were developed as described (Watkins et. al., (1998)). To isolate 10 individual clones, positive plaques from the initial screen were picked, replated at lower density (<103 phage/100-mm plate), and were screened by the same approach. Because the filters were probed with antigen at a concentration substantially below the Kd of the Fab 15 only variants displaying enhanced affinity were detectable. Multiple clones displaying higher affinities were identified following the screening of >106 variants from Hu II and >105 variants from the Hu III library using 82-mm filters containing 105 variants per filter (Table 20 I). Titration of the variants on immobilized CD40-Ig verified that multiple clones displayed affinities greater than the chimeric and humanized Fab (Fig. 2A, compare open squares, filled triangles with circles).

The framework/CDR mutations that conferred
25 enhanced affinity were identified by DNA sequencing.
Single-stranded DNA was isolated and the H and L chain
variable region genes were sequenced by the fluorescent
dideoxynucleotide termination method (Perkin-Elmer,
Foster City, CA). Unique variable region sequences were
30 identified in 10/13 Hu II variants and 4/5 Hu III
variants. Both the Hu II and Hu III variants contained
1-5 murine framework residues and 0-2 CDR3 mutations.
Representative examples are summarized in Table II. The
affinities of bacterially-expressed chimeric Fab and

63

certain variants from each of the libraries were characterized more thoroughly using surface plasmon resonance measurements to determine the association and dissociation rates of purified Fab with immobilized 5 CD40-Ig as described above.

Chimeric anti-CD40 had a dissociation constant  $K_d=48.3$  nM and, consistent with the screening results, the variants all displayed higher affinities with  $K_d$  ranging from 0.24 nM to 10.5 nM (Table II). Further optimization of LCDR3 of Hu III clone F4 resulted in the identification of a higher affinity ( $K_d=0.1$  nM) clone, L3.17, which contained a  $^{94}F\rightarrow Y$  mutation. The improved affinities of the anti-CD40 variants were predominantly the result of slower dissociation rates. However, the association rates of most variants were also enhanced, increasing by as much as  $\approx 3$ -fold (1.2 vs. 3.2 x  $10^6$  M<sup>-1</sup>s<sup>-1</sup> for chimeric anti-CD40 and clone L3.17, respectively).

Table II. Simultaneous optimization of framework and CDR residues.

Library	Clone	Kd (nM)	Murine Fr Residues*	CDR Mutations
	chimeri c	48.3	(43)	0
Hu I	19011	42.4	(2) H28, 48	0
	1H11	53.4	(4) H9, 28, 91, L49	0
	9A3	43.9	(3) H9, 28, 91	0
Hu II	CW43	10.53	(3) H9, 28, 91	HCDR3, 101A→R
	Y49K'	53.4	(4) H9, 28, 91, L49	HCDR3, 101A→R
	2B12	4.67	(5) H9, 28, 38, 46, 48	HCDR3, <sup>101</sup> A→K
Hu III	2B12	4.67	(5) H9, 28, 38, 46, 48	HCDR3, 101A→K
	288	2.81	(1) H28	HCDR3, 101A→K;
				LCDR3, 96R→Y
	F4	0.24	(1) H28	HCDR3, <sup>101</sup> A→K; LCDR3, <sup>96</sup> R→W
Hu IV	L3.17	0.10	(1) H28	HCDR3, <sup>101</sup> A→K LCDR3, <sup>94</sup> F→Y LCDR3, <sup>96</sup> R→W

\*The number of murine framework residues that differ from the most homologous human germline sequence based on 10 definition of CDRs of Kabat et. al. (1977, 1991) are indicated in parentheses. Differing murine framework residues retained in the humanized versions are located predominantly on the H chain (H) at the indicated positions. Hu I clone 1H11 and the CW43 derivative, 15 clone Y49K, contain a single differing L chain (L) framework residue at position 49.

<sup>†</sup>Clone Y49K was created by site-directed mutagenesis of clone CW43. The four clones within the shaded boxed region, 1H11, 9A3, CW43, and Y49K, were characterized to

65

demonstrate the co-operative interaction between L chain framework residue  ${\rm tyr^{49}}$  (human) and HCDR3 residue  ${\rm arg^{101}}$ .

The variants displaying enhanced affinity were tested for their ability to block the binding of gp39 5 ligand to the CD40 receptor. Immulon II microtiter plates were coated with 2  $\mu$ g/ml anti-murine CD8 to capture sgp39 fusion protein which expresses the CD8 domain. The plates were rinsed once with PBS containing 0.05% Tween 20, and were blocked with 3% BSA in PBS. The 10 plate was washed once with PBS containing 0.05% Tween 20 and was incubated with cell culture media containing saturating levels of sgp39 for 2 h at 25°C. Unbound sgp39 was aspirated and the plate was washed two times with PBS containing 0.05% Tween 20. Next, 25  $\mu$ l of 15 purified variant Fabs diluted serially 3-fold in PBS was added followed by 25  $\mu$ l of 4  $\mu$ g/ml CD40-human Iq in PBS. The plates were incubated 2 h at 25°C and were washed three times with PBS containing 0.05% Tween 20. Bound CD40-human Ig was detected following a 1 h incubation at 20 25°C with goat F(ab')<sub>2</sub> anti-human IgG Fcy-specific horseradish peroxidase conjugate (Jackson) diluted 10,000-fold in PBS. The plate was washed four times with PBS containing 0.05% Tween 20 and binding was quantitated colorimetrically by incubating with 1 mg/ml 25 o-phenylenediamine dihydrochloride and 0.003% hydrogen peroxide in 50 mM citric acid, 100 mM Na<sub>2</sub>HPO<sub>4</sub>, pH 5. The reaction was terminated by the addition of H2SO4 to a final concentration of 0.36 M and the absorbance at 490 nm was determined. Figure 2B shows purified variants 30 were tested for their ability to inhibit sgp39 binding to CD40-Ig. The ligand for the CD40 receptor, gp39, was captured in a microtiter plate and subsequently, varying amounts of purified chimeric (filled circles), Hu II-CW43 (open squares), Hu III-2B8 (filled triangles), Hu

WO 01/27160

II/III-2B12 (open triangles), and irrelevant (filled squares) Fab were co-incubated with 2  $\mu$ g/ml CD40-human Ig on the microtiter plate. The variants all inhibited the binding of soluble CD40-Ig fusion protein to immobilized 5 gp39 antigen in a dose-dependent manner that correlated with the affinity of the Fabs. For example, one of the most potent inhibitors of ligand binding to CD40-Ig fusion protein was variant 2B8, which was also one of the variants with the highest affinity for CD40. Variant 2B8 10 displayed ≈17-fold higher affinity for CD40 than did the chimeric Fab and inhibited ligand binding ≈7-fold more effectively.

66

PCT/US00/28435

Screening of the Hu I library identified two variants that were similar or identical in framework 15 sequence to the Hu II clone CW43 but displayed 5-fold lower affinities (Table II, clones 1H11 and 9A3). Clone 9A3 has an identical framework structure while 1H11 contained the murine lysine framework residue at L chain position 49. Sequence comparisons and site-directed 20 mutagenesis studies (data not shown) suggest that the beneficial arginine residue at HCDR3 position 101 might interact with L chain residue tyr49. To test this, L chain residue tyr49 of clone CW43 was mutated to the lysine murine framework residue, resulting in a variant 25 with a framework identical to clone 1H11 that also contained the beneficial arg<sup>101</sup> residue in HCDR3. The resulting mAb, termed Y49K, displayed 5-fold lower affinity than CW43. Thus, expression of tyrosine at L chain framework residue 49 or expression of arginine at 30 HCDR3 residue 101 alone had no beneficial effect on the mAb affinity, while the concomitant expression of tyrosine and arginine at these sites improved the mAb affinity 5-fold. The non-additive, or dependent nature of the mutations demonstrates that L chain residue tyr49

67

and HCDR3 residue arg<sup>101</sup> interact co-operatively to enhance the affinity of the mAb (Schreiber & Fersht, (1995)). In addition, the co-operative interaction that was observed between tyr<sup>49</sup> and arg<sup>101</sup> was also observed for variants that expressed lysine at HCDR3 position 101 (Table II).

Generally, interacting residues are spatially separated by no more than 7 Å (Schreiber & Fersht, 1995)). Figure 3 shows molecular modeling of anti-CD40 variant CW43. A top view of the anti-CD40 variant CW43 variable region structure was created by homology modeling to examine the spatial relationship of L chain framework residue Y49 and H chain CDR3 residue R101. The L chain is on the left and the H chain right with the H chain CDR3 loop depicted in red. The L chain framework residue 49 is in close proximity to the H chain CDR3 loop and is 7Å of the predicted interacting H chain CDR3 R101 residue. Although the interacting amino acids are located on distinct chains of the mAb, the residues are predicted to be within a range (7 Å) to permit co-operative interaction.

Throughout this application various publications have been referenced within parentheses.

The disclosures of these publications in their entireties are hereby incorporated by reference in this application in order to more fully describe the state of the art to which this invention pertains.

Although the invention has been described with reference to the disclosed embodiments, those skilled in the art will readily appreciate that the specific experiments detailed are only illustrative of the invention. It should be understood that various

68

modifications can be made without departing from the spirit of the invention. Accordingly, the invention is limited only by the following claims.

69

WO 01/27160 PCT/US00/28435

What is claimed is:

1. A method of conferring donor CDR binding affinity onto an antibody acceptor variable region framework, comprising:

- 5 (a) constructing a population of altered antibody variable region encoding nucleic acids, said population comprising encoding nucleic acids for an acceptor variable region framework containing a plurality of different amino acids at one or more acceptor

  10 framework region amino acid positions and donor CDRs containing a plurality of different amino acids at one or
  - (b) expressing said population of altered variable region encoding nucleic acids, and

more donor CDR amino acid positions;

- 15 (c) identifying one or more altered variable regions having binding affinity substantially the same or greater than the donor CDR variable region.
- The method of claim 1, wherein said one or more altered variable regions are identified by comparing
   the relative binding of said altered variable regions to said donor CDR variable region.
  - 3. The method of claim 1, wherein said one or more altered variable regions are identified by measuring the binding affinity of said altered variable regions.
- 4. The method of claim 1, wherein said one or more altered variable regions are identified by measuring the association rate (kon) or disassociation rate (koff) of said altered variable regions.

70

- 5. The method of claim 1, wherein said acceptor variable region framework is a heavy chain variable region framework.
- The method of claim 1, wherein said
   acceptor variable region framework is a light chain variable region framework.
- 7. The method of claim 1, wherein said framework amino acid positions are located in framework region 1, framework region 2, framework region 3 or 10 framework region 4.
  - 8. The method of claim 1, wherein said donor CDR amino acid positions is located in CDR1, CDR2 or CDR3.
- 9. The method of claim 1, wherein said one or 15 more amino acid positions in said acceptor framework region is selected by differences in amino acid identity between corresponding positions in donor and acceptor framework regions.
- 20 10. The method of claim 1, wherein said one or more amino acid positions in said acceptor framework region is selected as being a canonical framework residue.
- 11. The method of claim 1, wherein said one or 25 more amino acid positions in said acceptor framework region is selected as being exposed to solvent.
  - 12. The method of claim 1, wherein said one or more amino acid positions in said acceptor framework region is selected by a characteristic within the group

consisting of being proximal to a CDR, predicted to contact the opposite domain in the  $V_1-V_H$  interface, lack of relatedness to the donor framework amino acid position at that position and predicted to modulate CDR activity.

71

- 5 13. The method of claim 1, wherein said one or more amino acid positions in said donor CDR isselected as being a CDR residue as defined by Kabat
- 14. The method of claim 5, wherein said altered variable regions are coexpressed with a light10 chain variable region.
  - 15. The method of claim 6, wherein said altered variable region is coexpressed with a heavy chain variable region.
- 16. A method of simultaneously grafting and 15 optimizing the binding affinity of a variable region binding fragment, comprising:
- (a) constructing a population of altered heavy chain variable region encoding nucleic acids comprising an acceptor variable region framework containing donor
   CDRs and a plurality of different amino acids at one or more framework region and CDR amino acid positions;
- (b) constructing a population of altered light chain variable region encoding nucleic acids comprising an acceptor variable region framework containing donor
   CDRs and a plurality of different amino acids at one or more framework regions and CDR amino acid positions;
- (c) coexpressing said populations of heavy and light chain variable region encoding nucleic acids to produce diverse combinations of heteromeric variable
   region binding fragments, and

WO 01/27160

(d) identifying one or more heteromeric variable region binding fragments having affinity substantially the same or greater than the donor CDR heteromeric variable region binding fragment.

72

PCT/US00/28435

- 5 17. The method of claim 16, wherein said one or more heteromeric variable region binding fragments are identified by comparing the relative binding of said heteromeric variable region binding fragments to said donor CDR heteromeric variable region binding fragment.
- 10 18 The method of claim 16 wherein said one or more heteromeric variable region binding fragments are identified by measuring the binding affinity of said heteromeric variable region binding fragments.
- 19. The method of claim 16, wherein said one
  15 or more heteromeric variable region binding fragments are
  identified by measuring the association rate (kon) or
  disassociation rate (koff) of said heteromeric variable
  region binding fragments.
- 20. The method of claim 16, wherein said
  20 framework amino acid positions are located in framework
  region 1, framework region 2, framework region 3 or
  framework region 4.
- 21. The method of claim 16, wherein said donor CDR amino acid positions is located in CDR1, CDR2 or 25 CDR3.

- 22. The method of claim 16, wherein said one or more amino acid positions in said acceptor framework region is selected by differences in amino acid identity between corresponding positions in donor and acceptor framework regions.
- 23. The method of claim 16, wherein said one or more amino acid positions in said acceptor framework region is selected as being a canonical framework 10 residue.
  - 24. The method of claim 16, wherein said one or more amino acid positions in said acceptor framework region is selected as being exposed to solvent.
- 25. The method of claim 16, wherein said one or more amino acid positions in said acceptor framework region is selected by a characteristic within the group consisting of being proximal to a CDR, predicted to contact the opposite domain in the  $V_L-V_R$  interface, lack of relatedness to the donor framework amino acid position at that position and predicted to modulate CDR activity.
  - 26. The method of claim 16, wherein said one or more amino acid positions in said donor CDR is selected as being a CDR residue as defined by Kabat.
- 27. A method of optimizing the binding25 affinity of an antibody variable region, comprising:
- (a) constructing a population of antibody variable region encoding nucleic acids from a parent variable region encoding nucleic acid, said population comprising two or more CDRs containing a plurality of different amino acids at one or more CDR amino acid positions;

WO 01/27160

74

PCT/US00/28435

- (b) expressing said population of variable region encoding nucleic acids, and
- (c) identifying one or more variable regions having binding affinity substantially the same or greater 5 than the parent variable region.
  - 28. The method of claim 27, wherein said one or more variable regions are identified by comparing the relative binding of said variable regions to said parent variable region.
- 29. The method of claim 27, wherein said one or more variable regions are identified by measuring the binding affinity of said variable regions.
- 30. The method of claim 27, wherein said one or more variable regions are identified by measuring the association rate (kon) or disassociation rate (koff) of said variable regions.
  - 31. The method of claim 27, wherein said variable region is a heavy chain variable region.
- 32. The method of claim 27, wherein said 20 variable region is a light chain variable region.
  - 33. The method of claim 27, wherein said two or more CDRs are selected from the group consisting of CDR1, CDR2 or CDR3.
- 34. The method of claim 27, wherein said one or more amino acid positions in said two or more CDRs is selected as being a CDR residue as defined by Kabat.

WO 01/27160

PCT/US00/28435

- 35. The method of claim 27, wherein said variable regions are coexpressed with a light chain variable region.
- 36. The method of claim 32, wherein said 5 variable regions are coexpressed with a heavy chain variable region.
- 37. The method of claim 27, wherein said antibody variable region is selected from the group consisting of native, grafted, altered and optimized variable regions.
  - 38. A method of optimizing the activity of a catalytic antibody variable region, comprising:
- (a) constructing a population of heavy chain variable region encoding nucleic acids from a parent
   15 heavy chain variable region encoding nucleic acid, said population comprising two or more CDRs containing a plurality of different amino acids at one or more CDR amino acid positions;
- (b) constructing a population of light chain 20 variable region encoding nucleic acids from a parent light chain variable region encoding nucleic acid, said population comprising two or more CDRs containing a plurality of different amino acids at one or more CDR amino acid positions;
- 25 (c) coexpressing said population of heavy and light chain variable region encoding nucleic acids containing said two or more CDRs having said plurality of different amino acids at one or more CDR positions to produce diverse combinations of heteromeric variable 30 region catalytic fragments, and
  - (d) identifying one or more heteromeric variable regions having optimized catalytic activity

76

compared to said parent catalytic antibody variable region.

- 39. The method of claim 38, wherein said one or more heteromeric variable regions are identified by comparing the relative catalytic activity of said heteromeric variable regions to said parent variable region.
- 40. The method of claim 38, wherein said one or more heteromeric variable regions are identified by 10 measuring a substrate association rate (kon), a substrate disassociation rate (koff), substrate binding affinity, a transition state binding affinity, a turnover rate or a Km.
- 41. The method of claim 38, wherein said two or more CDRs are selected from the group consisting of CDR1, CDR2 and CDR3.
  - 42. The method of claim 38, wherein said one or more amino acid positions in said two or more CDRs is selected as being a CDR residue as defined by Kabat.



## **V<sub>H</sub> Domain**

CD40 10 20 30 40
QIQLVQSGPELKKPGETVRISCKASGYAFTTTGMQWVQEMPGKGLKWIG
VH7 QVQLVQSGSELKKPGASVKVSCKASGYTFTSYAMNWVRQAPGQGLEWMG

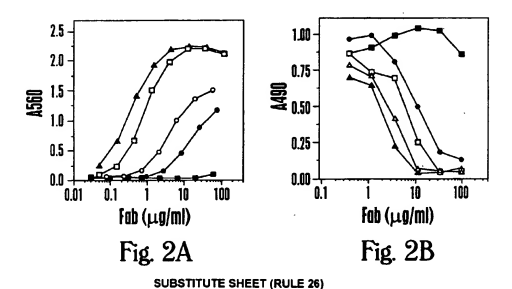
CD40 50 60 70 80 abc 90 WINTHSGVPKYVEDFKGRFAFSLETSANTAYLQISNLKNEDTATYFCVR

VH7 <u>WINTNTGNPTYAQGFTG</u>RFVFSLDTSVSTAYLQISSLKAEDTAVYYCAR

abcde 110
SGNGNYDLAYFAYWGQGTLVTVSA

JH4 YFDYWGQGTLVTVSS

Fig. 1



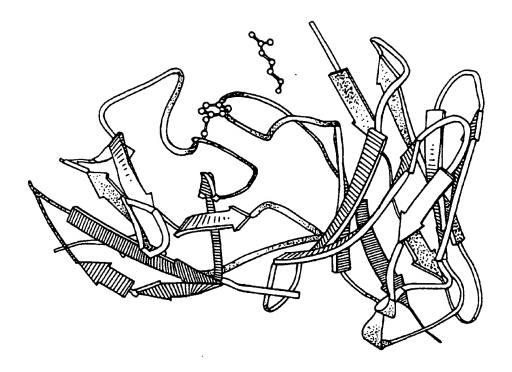
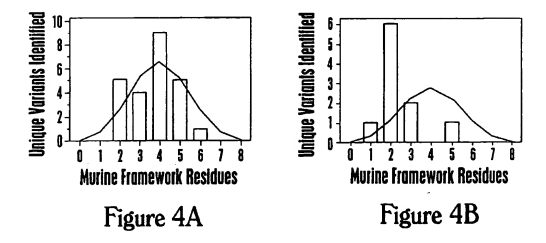


Figure 3



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